# UK Patent Application (19) GB (11) 2 264 496 (13) A

(43) Date of A publication 01.09.1993

- (21) Application No 9302411.5
- (22) Date of filing 08.02.1993
- (30) Priority data (31) 841114
- (32) 25.02.1992

(33) US

- (71) Applicant **United States Department of Energy** 1000 Independence Avenue S.W, Washington D.C 20585, District of Columbia, **United States of America**
- (72) Inventors Richard Alan Keller Mark Lewis Hammond **Babetta Louise Marrone** John Calvin Martin James Hubert Jett
- (74) Agent and/or Address for Service Potts, Kerr & Co 15 Hamilton Square, Birkenhead, Merseyslde, L41 6BR, United Kingdom

- (51) INT CL<sup>5</sup> G01N 21/64, C12Q 1/68
- (52) UK CL (Edition L) C3H HB3 G1B BAC BBN B121
- (56) Documents cited WO 90/10718 A WO 89/03432 A
- (58) Field of search UK CL (Edition L) C3H HB3, G1B BAC BAX BBN INT CL<sup>5</sup> C12Q, G01N Online database: WPI

#### (54) Sizing DNA by induced fluorescence

(57) A method is provided for obtaining DNA fingerprints using high speed detection systems, such as flow cytometry to determine unique characteristics of DNA pieces from a selected sample. In one characterization the DNA piece is fragmented at preselected sites to produce a plurality of DNA fragments. The DNA piece or the resulting DNA fragments are treated with a dye effective to stain stoichiometrically the DNA fragments. The fluorescence from the dye in the stained fragments is then examined to generate an output functionally related to the number of nucleotides in each one of the DNA fragments. The intensity of the fluorescence emissions from each fragment is directly proportional to the fragment length. Additional dyes can be bound to the DNA piece and DNA fragments to provide information additional to length information. Oligonucleotide specific dyes and/or hybridization probes can be bound to the DNA fragments to provide information on oligonucleotide distribution or probe hybridization to DNA fragments of different sizes.

### BACKGROUND OF THE INVENTION

This invention relates to DNA analysis and, more particularly, to DNA fragment size distribution analysis and sorting. This invention is the result of a contract with the Department of Energy (Contract No. W-7405-ENG-36).

The human genome is comprised of some three billion nucleotides forming the 22 pairs of chromosomes plus 2 autosomes, each with continuous DNA pieces of 50-500 The organization and sequence of DNA million nucleotides. forming the human genome contains unique information about the source that provides the DNA. One method for accessing this information is to fragment the DNA at sites with known characteristics and then to analyze the distribution of fragment sizes, i.e., the number of nucleotides in each fragment between each of the sites. Polymorphisms in the genome structure lead to substantial variation in the fragment sizes obtained from fragmentation of DNA pieces and allow one to differentiate one person from another or to form a basis for assessing a person's susceptibility to genetic diseases. Analysis of these polymorphisms is often referred to as DNA fingerprinting.

fingerprinting is an important medical diagnostic forensic applications to additional with tool, identification, medical genetics, monitoring the effects of basic molecular biology and environmental mutagens, One form of DNA fingerprinting involves research. "restriction fragment length polymorphism" (RFLP) restriction enzymes are used to cut a DNA piece from a specific source into shorter pieces, or fragments, of DNA. RFLP provides a unique pattern of DNA fragments containing a unique DNA sequence ordered by fragment size (the DNA specimen is digested with DNA a fingerprint) when There are many known restriction restriction enzymes. enzymes and each recognizes a specific DNA sequence of four to twelve base pairs at which it cuts the DNA, resulting in smaller fragments of DNA.

Once the DNA piece has been cut into many fragments, electrophoresis is conventionally used to separate the An electric field is placed across a fragments by size. gel containing the fragments causing the smaller fragments to move faster than the larger ones. Gel electrophoresis is a well known technique and has been used to produce band patterns of DNA fragments that form a fingerprint to identify the individual source of the DNA piece under The band patterns of specific DNA sequences are analysis. conventionally visualized by binding radioactive DNA probes to the separated DNA fragments and exposing suitable film See, e.g., J.I. to the radioactive labeled fragments. Thornton, "DNA Profiling," C&EN, pp. 18-30 (November 20, 1989); K. Heine, "DNA on Trial," Outlook 26:4, pp. 8-14 In one variation, the fragment ends are tagged (1989).with a fluorescent dye so that the fragment migration time along a known path length in an electrophoretic gel can be determined by automated fluorescence detection. See, e.g., A.V. Carrano, "A High-Resolution, Fluorescence-Based, Semiautomated Method for DNA Fingerprinting," 4 Genomics, pp. 129-136 (1989).

There are, however, several limitations on the use of electrophoresis, particularly where large fragment gel sizes and radioactive labeling are involved. instances, the electrophoretic separation process takes considerable time to provide resolution for large size The development of images from radioactive fragments. probes is an additional time consuming step and has health associated concerns environmental hazards and Additionally, the distribution of radioactive materials. fragment sizes is logarithmic so that the separation, i.e., resolution, between large fragments is less than for small Electrophoresis also requires relatively large fragments. amounts of DNA to obtain a recognizable pattern.

It is desirable to provide a DNA fragment size analysis technique that uses only small quantities of DNA (maybe only a single strand), provides size information within a short time, and has a high resolution between fragment These and other problems of the prior art are sizes. wherein invention present the addressed by obtain to a used cytometry-based techniques are distribution of DNA fragment sizes from a DNA piece.

Accordingly, it is an object of the present invention to provide rapid determination of DNA fragment sizes.

It is another object of the present invention to obtain a high resolution of DNA fragments, particularly long fragments.

One other object of the present invention is to require only a small DNA sample to provide accurate DNA fragment size fingerprints.

Yet another object of the present invention is to enable fragment length detection without the use of radioactive labels.

A further object of the present invention is to use fluorescent intensities to determine the length of DNA fragments.

still another object of the present invention is to use the sorting capabilities associated with flow cytometry to sort the fragments by size, i.e., length, for further study.

Additional objects, advantages and novel features of the invention will be set forth in part in the description which follows, and in part will become apparent to those skilled in the art upon examination of the following or may be learned by practice of the invention. The objects and advantages of the invention may be realized and attained by means of the instrumentalities and combinations particularly pointed out in the appended claims.

#### SUMMARY OF THE INVENTION

To achieve the foregoing and other objects, and in accordance with the purposes of the present invention, as embodied and broadly described herein, the method of this invention may comprise the use of an induced fluorescence to quantitate the length of DNA fragments. A piece of DNA is fragmented at preselected sites to produce a plurality of DNA fragments. All of the DNA fragments are treated with a dye effective to stoichiometrically stain the nucleotides along the DNA fragments. The stained DNA

fragments are then fluorescently examined to generate an output functionally related to the number of nucleotides in each one of the DNA fragments. In one output, the intensity of the fluorescence emissions from each fragment is directly proportional to the fragment length.

## DETAILED DESCRIPTION OF THE INVENTION

In accordance with the present invention, DNA polymorphisms are characterized, i.e., "fingerprinted," using flow cytometry-based techniques to provide a rapid analysis of DNA fragment sizes obtained by fragmenting a selected DNA piece with one or more enzymes selected to cleave DNA at known sequence sites. One exemplary procedure for sizing is the following:

- 1. A DNA piece from a selected source is fragmented by enzyme digestion to provide a solution of DNA fragments. The nucleotides comprising the DNA piece may be stained, i.e., labeled, with an appropriate fluorescent dye either before or after the DNA piece is fragmented.
- 2. The stained DNA fragments are passed through a detection apparatus at a concentration and rate effective to provide only one fragment in the fluorescence excitation volume at any one time.
- 3. Each stained DNA fragment is excited, e.g., with laser irradiation, in the excitation volume and the resulting fluorescence intensity is measured, wherein the intensity of the induced fluorescence is a measure of the amount of stain on the fragment and concomitant fragment length.
- 4. The number of fragments at each different intensity provides an analysis of the number of fragments of each length produced from the DNA piece by the selected enzyme or enzymes.

A DNA piece may be first selected from any suitable source, e.g., blood, tissue samples, semen, laboratory research specimens, etc. The DNA piece is then fragmented using an enzyme chosen for a particular application of the analysis. One particularly useful type of enzyme is a restriction endonuclease that recognizes specific sites, i.e., specific nucleotide sequences, and cleaves the DNA piece within the identified sequence. For example, the enzyme Eco RI cuts at the double piece recognition site

#### ···GAATTC···

#### ···CTTAAG···

Hundreds of different restriction enzymes and their respective cleavage sites are known. It will be appreciated that identical DNA pieces from a single source might be digested with different enzymes to yield a family of fingerprints. Alternatively, a DNA piece may be digested with multiple enzymes to further particularize the fragment size distribution analysis.

Fragmentation, i.e., digestion, of a DNA piece with a selected enzyme is a well-known process, where the optimum the specified enzyme by digestion conditions are A generic restriction enzyme process for use manufacturer. with 0.2-1  $\mu$ g of DNA is given by J. Sambrook et al., 5.28-5.33, Cold Spring Harbor Cloning, pp. Molecular Laboratory Press (1989):

- 1. Place the DNA solution in a sterile microfuge tube and mix with sufficient water to give a volume of 18  $\mu L$ .
- 2. Add 2  $\mu L$  of an appropriate restriction enzyme digestion buffer and mix by tapping the tube. An exemplary buffer may be formed as follows:

200 mM potassium glutamate

- 50 mM Tris-acetate (pH 7.5)
- 20 mM magnesium acetate
- 100  $\mu$ g/mL bovine serum albumin (Fraction V; Sigma)
  - 1 mM  $\beta$ -mercaptoethanol.
- 3. Add 1-2 units of restriction enzyme and mix by tapping the tube, where 1 unit of enzyme is defined as the amount required to digest 1  $\mu g$  of DNA to completion in 1 hour in the recommended buffer and at the recommended temperature in a 20- $\mu L$  reaction.
- 4. Incubate the mixture at the appropriate temperature for the required period of time.
- 5. Stop the reaction by adding 0.5 M EDTA (pH 8.0) to a final concentration of 10 mM.

DNA can be fragmented by a variety of other techniques in addition to restriction enzyme digest:

- 1. DNase hypersensitivity sites (DNase footprinting). Chromatin digestion by DNase will produce fragments of various lengths due to differences in proteins that bind to the DNA and prevent cutting of the DNA by a DNase at sites where protein is bound (D.J. Galas et al., Nucleic Acids Res., 5:3157 (1987)).
- fluorescent RNA probe is synthesized complementary to a normal, or wild type, DNA sequence of interest. This complementary probe is then annealed to the target DNA that is to be analyzed. To determine if a single nucleotide mismatch exists between the fluorescent probe strand and the target DNA strand, the RNA:DNA hybrid is treated with RNase A. RNase A specifically cleaves single stranded regions of RNA, thus cleaving the single base pair mismatch region in the fluorescent RNA strand of the RNA:DNA hybrid. (See R.M. Myers et al., Science 230:1242 (1985) and E.F. Winter et al., Proc. Natl. Acad. Sci. 82:7525 (1985)).

- RecA-assisted restriction endonuclease cleavage. 3. oligonucleotides coated with RecA protein are Short annealed to the complementary target DNA sequence. DNA:oligonucleotide hybrid is treated with Eco RI methylase Eco RI sites that are not protected by the enzyme. methylated while oligonucleotide oligonucleotide are sites remain unaffected. Eco RI RI protected Eco restriction endonuclease will cleave only at protected sites (i.e., unmethylated). This method has been used to base pairs. (L.J. Ferrin, generate fragments>500,000 Science, 254:1494 (1991).
- 4. DNA fragmentation can also be accomplished by techniques other than enzyme digestion. For example, ultrasonic excitation at different frequencies might be used to produce a family of size distributions. Various chemicals also react with the nucleotides and may be used to fragment DNA pieces.

The DNA fragments must be stained with a fluorescent dye for flow cytometric analysis. A fluorescent dye is selected to bind stoichiometrically to the DNA fragments. The complex may be formed in different ways, i.e., single stranded DNA, double stranded DNA, specific base pairs, Well known dyes include ethidium bromide, acridine propidium iodide, DAPI, Hoechst, chromomycin, orange, mithramycin, 9 amino acridine, ethidium bromide heterodyne, asymmetric cyanine dyes, or combinations of these dyes for The selected dye or dyes bind to the energy transfer. oligonucleotides stoichiometrically along the DNA sequence, i.e., the binding sites along the fragments are such that the total number of dye molecules along any length of DNA is proportional to the number of base pairs (bp's) forming For example, under the staining protocol set the DNA.

forth below, the number of ethidium bromide molecules bound to a DNA fragment is stoichiometric and can be as high as one-half the number of bp's. See, e.g., C.R. Cantor et al., "Binding of Smaller Molecules to Nucleic Acids," in Biophysical Chemistry, Part III: The Behavior of Biological Macromolecules, p. 1251, W.H. Freeman and Company, 1980.

One exemplary procedure for staining with ethidium bromide is:

Add DNA sample to a solution containing 1-5  $\mu g$  of ethidium bromide per mL of solution and TE 8.0 buffer. A suitable buffer is available from GIBCO and is 10 mM Tris-HCL and 1 mM EDTA, pH 8.0. The reaction is complete in 5-10 minutes at room temperature.

The stained DNA fragments are now analyzed using sensitive fluorescence detection techniques to determine the fluorescence intensity from each fragment passing through a detection region and having a high resolution to distinguish adjacent fragment sizes. Theoretically, while only a single DNA piece is needed to obtain the desired distribution analysis, a typical solution will be formed from many DNA pieces and a relative DNA fragment size distribution is obtained. The DNA fragments will typically range in size from 100 bp to 500,000 bp.

It is well-known how to form a sequential flow stream of particles for use in a flow cytometer or similar sensitive fluorescence detection apparatus. See, e.g., U.S. Patent 3,710,933, issued January 16, 1973, to Fulwyler et al. and <u>Flow Cytometry and Sorting</u>, 2nd Ed., ed. M.R. Melamed et al., Wiley-Liss, New York, 1990, incorporated herein by reference. A dilute solution of the DNA fragments is formed to a low concentration effective to provide the fragments spaced apart in the flow stream so

that only a single fragment is present in the excitation The solution of DNA fragments is then injected volume. within a laminar sheath flow stream for passage through the detection chamber for laser excitation of one fragment at a The flow rates of the sample and the sheath are time. maintain separation between particles and to adjusted to optimum time for each particle in the the provide An optimum time is determined from a excitation source. consideration of sizing rate, detection sensitivity, and photostability of the dye tags. A suitable excitation source is selected to initiate fluoresce in the dye used to stain the DNA fragments. For example, an argon laser at 488 nm is effective to cause ethidium bromide to fluoresce in a band around 600 nm.

The sensitivity of conventional flow cytometry system is improved by providing a small excitation volume, e.g.,  $10-20~\mu\text{m}$  diameter and  $100~\mu\text{m}$  length, with a tightly focused laser beam. See, e.g., J.H. Hahn et al., "Laser-Induced Fluorescence Detection of Rhodamine-6G at  $6\times10^{-15}\text{M}$ ," Appl. Spectrosc. 45:743~(1991), describing a probe volume of 11 pL, incorporated herein by reference. The small probe volume greatly reduces the amount of background emission, i.e., noise, in the output signal.

The laser excitation may be a pulsed laser with a pulse of e.g., about 70ps full width, with time gating to differentiate between dye emission photons (delayed) and Raman scattering photons (prompt). See, e.g., E.B. Shera et al., "Detection of Single Fluorescent Molecules," Chem. Phys. Lett. 174:553 (November 1990), The prompt scattered photons occur within the laser pulse time while the dye emissions decay with a several nanosecond lifetime so that a delayed window is effective for discrimination of

scatter photons. Raman from photons fluorescence Alternatively, the laser may be a cw laser. See, e.g., S.A. Soper, "Single-Molecule Detection of Rhodamine-6G in Continuous Laser Wave Using Solutions Ethanolic The number of Excitation, " Anal. Chem. 63:432 (1991). emitted photons can be increased by increasing the DNA fragment transit time through the laser beam and by selecting a dye and solvent with high photostability for The number of detected photons (photoelectrons) the dye. is also increased by increasing the sensitivity of the Furthermore, the present invention detection apparatus. involves DNA fragments rather than single molecules so that fragment length so that a larger output longer the fluorescence intensity is obtained.

It will also be appreciated that the solution may contain some dye that was not bound to the DNA fragments. This dye will be excited along with bound dye and the system must discriminate between the fluorescence from the unbound and the bound dye. In one embodiment, a pulsed and gated detection technique may be used to provide For example, the excited state discrimination. this lifetimes for the unbound and bound ethidium bromide are 2 ns and 23 ns, respectively. Thus, the detection system can be gated to detect only the fluorescence from the bound ethidium bromide and, hence, provide an output signal functionally related to the length of the DNA fragment. Alternatively, a dye might be selected that provides different fluorescence or absorption wavelengths in the For example, a series of bound and unbound states. asymmetric cyanine dyes are reported by I.D. Johnson et al., "Asymmetric Cyanine Dyes for Fluorescent Staining and Fluorescence Acids," Nucleic of Quantification

Spectroscopy, Abstract 1806, FASEB J. 6:A314, No. 1 (January 1992).

Polarized fluorescence emission also provides a means from unbound dye molecules. discriminating bound of Fluorescence polarization of unbound DNA dyes is  $\leq 0.05$ , fluorescence polarization of DNA bound the whereas fluorochromes can be between 0.20 and 0.30. See, e.g., L.S. Cram et al., "Fluorescence Polarization and Pulse Analysis of Chromosomes by a Flow System," J. Width Histochem. Cytochem. 27:445, No. 1 (1979); T.M. Jovin, "Fluorescence Polarization and Energy Transfer: Theory and Application," Flow Cytometry and Sorting, Ed. M.R. Melamed et al., pp. 156, John Wiley & Sons (1979). Discrimination is accomplished by using a polarized excitation source and detecting the emissions through a polarization filter a fluorescence detector. The of front placed in polarization filter is aligned with the polarization direction parallel to the polarization direction of the excitation source.

an energy excitation, laser continuous CW For transfer-type scheme may be used to distinguish bound and If a second dye is also bound to unbound dye molecules. the DNA, the bound first dye molecules used for fragment sizing will be in a close proximity to the second dye molecules so that excitation of the second dye molecules result in energy transfer from the second dye will molecules to the first dye molecules. The unbound first and second dye molecules in the surrounding fluid will not be in proximity effective for energy transfer. Thus, only the bound first dye molecules will fluoresce for fragment length determination when the second dye molecules are excited.

Spectroscopy, Abstract 1806, FASEB J. 6:A314, No. 1 (January 1992).

Polarized fluorescence emission also provides a means from unbound dye molecules. discriminating bound Fluorescence polarization of unbound DNA dyes is  $\leq 0.05$ , of DNA bound fluorescence polarization the whereas fluorochromes can be between 0.20 and 0.30. See, e.g., L.S. Cram et al., "Fluorescence Polarization and Pulse Analysis of Chromosomes by a Flow System," J. Width Histochem. Cytochem. 27:445, No. 1 (1979); T.M. Jovin, "Fluorescence Polarization and Energy Transfer: Theory and Application," Flow Cytometry and Sorting, Ed. M.R. Melamed et al., pp. 156, John Wiley & Sons (1979). Discrimination is accomplished by using a polarized excitation source and detecting the emissions through a polarization filter a fluorescence detector. The of placed in front polarization filter is aligned with the polarization direction parallel to the polarization direction of the excitation source.

energy excitation, an laser continuous CW For be used to distinguish bound and transfer-type scheme may If a second dye is also bound to unbound dye molecules. the DNA, the bound first dye molecules used for fragment sizing will be in a close proximity to the second dye molecules so that excitation of the second dye molecules will result in energy transfer from the second dye molecules to the first dye molecules. The unbound first and second dye molecules in the surrounding fluid will not be in proximity effective for energy transfer. Thus, only the bound first dye molecules will fluoresce for fragment length determination when the second dye molecules are excited.

resolution improves as  $N^{1/2}$ . For N=1000,  $\sigma=3.87$  and R=0.0775%. Sizing 10, 100, or even 10000 identical fragments is not a problem. There are many more fragments than 10000 in a typical electrophoresis band. Thus, it can be seen that the resolution can be much better than 1% on a 100,000 bp fragment, whereas a resolution of only 10-20% would be expected for separation of fragments in the 100,000 bp range by gel electrophoresis and the resolution degrades further as fragment length increases.

DNA fingerprinting according to the present invention can also be done very rapidly. A typical DNA fingerprint by electrophoresis has about 50 bands. At 1000 fragments per band, 50 bands would require only about 8.3 minutes to develop a fingerprint at a fragment analysis rate of 100 fragments/second. If only a single band is required for the desired resolution, the analysis time would be only about 1 second. If the desired resolution requires 100 fragments per band, then the 50 band analysis would take only about 50 seconds.

present invention, the adaptation of In an hybridization probes can be bound to the DNA restriction associated with fragment length sizes. fragments and Hybridization probes are conventionally formed containing a probe dye and hybridized to DNA fragments formed by base matching from the DNA piece being investigated. pair Excitation of the hybridized DNA fragments could then be designed to excite both the size-measuring dye and the probe dye so that correlation of the fluorescent outputs would associate the probe with various fragment lengths. In situ probe hybridization to DNA is discussed in Methods in Cell Biology, Vol. 33, Z. Darzynkiewicz et al. Ed., York 1990), Chapter 37, (New Inc. Press, Academic

resolution improves as  $N^{1/2}$ . For N=1000,  $\sigma$ =3.87 and R=0.0775%. Sizing 10, 100, or even 10000 identical fragments is not a problem. There are many more fragments than 10000 in a typical electrophoresis band. Thus, it can be seen that the resolution can be much better than 1% on a 100,000 bp fragment, whereas a resolution of only 10-20% would be expected for separation of fragments in the 100,000 bp range by gel electrophoresis and the resolution degrades further as fragment length increases.

DNA fingerprinting according to the present invention can also be done very rapidly. A typical DNA fingerprint by electrophoresis has about 50 bands. At 1000 fragments per band, 50 bands would require only about 8.3 minutes to develop a fingerprint at a fragment analysis rate of 100 fragments/second. If only a single band is required for the desired resolution, the analysis time would be only about 1 second. If the desired resolution requires 100 fragments per band, then the 50 band analysis would take only about 50 seconds.

invention, present the of adaptation an In hybridization probes can be bound to the DNA restriction associated with fragment length sizes. fragments and Hybridization probes are conventionally formed containing a probe dye and hybridized to DNA fragments formed by base matching from the DNA piece being investigated. pair Excitation of the hybridized DNA fragments could then be designed to excite both the size-measuring dye and the probe dye so that correlation of the fluorescent outputs would associate the probe with various fragment lengths. In situ probe hybridization to DNA is discussed in Methods in Cell Biology, Vol. 33, Z. Darzynkiewicz et al. Ed., (New York 1990), Chapter 37, Press, Inc. Academic

Conventional sorting apparatus, as discussed in U.S. Patent 3,710,933 and in T. Lindmo, "Flow Sorters for Biological Cells," Flow Cytometry and Sorting, Second Edition, Ed. M. Melamed et al., pp. 145-169, John Wiley & Sons (1990), uses the fluorescence output signals discussed After the fragments have passed through the above. generating the output, the for volume excitation hydrodynamic flow stream is broken into droplets by, e.g., ultrasonic vibrations, where each drop contains no more Drops containing DNA fragments that than one fragment. have a selected fluorescence response to an excitation are charged by the application of a high voltage pulse across which then pass through charged plates that the drops, generate an electrostatic field to selectively deflect the charged drops. The charge applied to the selected drops is controlled by circuitry that is responsive to fluorescent from the excitation volume within the flow emissions cytometer, where the charging pulse is activated to produce deflection of drops containing a material emitting fluorescence at a selected wavelength and intensity.

While the above description has been directed to DNA pieces, the process is equally applicable to RNA strands. Any reference to DNA in this case should be construed to include RNA. Likewise, the form of signal detected is taught to be fluorescence. However, any form of light emission may be obtained, depending on the specific dye, such that the term fluorescence should be interpreted to include phosphorescence and luminescence. Further, the DNA or RNA being fingerprinted may not necessarily be from humans, since all organisms have a genome that determines their specific characteristics.

The foregoing description of the preferred embodiments of the invention have been presented for purposes of It is not intended to be illustration and description. exhaustive or to limit the invention to the precise form disclosed, and obviously many modifications and variations in light of the above teaching. The possible are embodiments were chosen and described in order to best explain the principles of the invention and its practical application to thereby enable others skilled in the art to best utilize the invention in various embodiments and with various modifications as are suited to the particular use It is intended that the scope of the contemplated. invention be defined by the claims appended hereto.

#### WHAT IS CLAIMED IS:

1. A method for sizing DNA fragments, each containing some number of nucleotides, by induced fluorescence, comprising the steps of:

fragmenting a piece of DNA at preselected sites within said piece of DNA to produce a plurality of DNA fragments;

staining said piece of DNA or said DNA fragments with a first dye effective to stain stoichiometrically said nucleotides; and

fluorescently examining said DNA fragments after staining to generate a fluorescence intensity signal from said first dye that is functionally related to the number of nucleotides in each one of said DNA fragments.

- 2. A method according to Claim 1, wherein the step of fragmenting said piece of DNA comprises the step of subjecting said piece to a restriction enzyme selected to cleave said piece at a known combination of base pairs.
- 3. A method according to Claim 1, wherein the step of fragmenting said piece of DNA includes the steps of:

binding a selected DNA sequence to said DNA piece to identify selected sites along said piece; and

cleaving said DNA piece at said selected sites to form said plurality of DNA fragments.

- 4. A method according to Claim 1, wherein the step of staining said DNA piece or said DNA fragments with a first dye includes the step of adding said DNA piece or said DNA fragments to a solution containing a dye selected from the group consisting of ethidium bromide, acridine orange, propidium iodide, DAPI, Hoechst chromomycin, mithramycin, 9 amino acridine, and ethidium acridine heterodyne.
- 5. A method according to Claim 2, wherein the step of staining said DNA piece of said DNA fragments with said first dye includes the step of adding said DNA fragments to a solution containing a dye selected from the group consisting of ethidium bromide, acridine orange, propidium iodide, DAPI, Hoechst, chromomycin, mithramycin, 9 amino acridine, and ethidium acridine heterodyne.
- 6. A method according to Claim 3, wherein the step of staining said DNA piece or said DNA fragments with said first dye includes the step of adding said DNA fragments to a solution containing a dye selected from the group consisting of ethidium bromide, acridine orange, propidium iodide, DAPI, Hoechst, chromomycin, mithramycin, 9 amino acridine, and ethidium acridine heterodyne.
- 7. A method according to Claim 1, wherein the step of fluorescently examining said DNA fragments includes the steps of:

forming a flow stream of said DNA fragments serially spaced apart in said flow stream;

illuminating said flow stream with a laser effective to cause said first dye to fluoresce; and

measuring the intensity of the total fluorescence from each one of said fragments of DNA in said flow stream.

- 8. A method according to Claim 7, wherein said step of measuring said total fluorescence intensity further includes the step of differentiating dye molecules bound to said DNA fragments from unbound dye molecules.
- 9. A method according to Claim 2, wherein the step of fluorescently examining said DNA fragments includes the steps of:

forming a flow stream of said DNA fragments serially spaced apart in said flow stream;

illuminating said flow stream with a laser effective to cause said first dye to fluoresce; and

measuring the intensity of the total fluorescence from each one of said fragments of DNA in said flow stream.

- 10. A method according to Claim 9, wherein said step of measuring said total fluorescence intensity further includes the step of differentiating dye molecules bound to said DNA fragments from unbound dye molecules.
- 11. A method according to Claim 3, wherein the step of fluorescently examining said DNA fragments includes the steps of:

forming a flow stream of said DNA fragments serially spaced apart in said flow stream;

illuminating said flow stream with a laser effective to cause said first dye to fluoresce; and

measuring the intensity of the total fluorescence from each one of said fragments of DNA in said flow stream.

12. A method according to Claim 11, wherein said step of measuring said total fluorescence intensity further includes the step of differentiating dye molecules bound to said DNA fragments from unbound dye molecules.

13. A method according to Claim 4, wherein the step of fluorescently examining said DNA fragments includes the steps of:

forming a flow stream of said DNA fragments serially spaced apart in said flow stream;

illuminating said flow stream with a laser effective to cause said first dye to fluoresce; and

measuring the intensity of the total fluorescence from each one of said fragments of DNA in said flow stream.

- 14. A method according to Claim 13, wherein said step of measuring said total fluorescence intensity further includes the step of differentiating dye molecules bound to said DNA fragments from unbound dye molecules.
- 15. A method according to Claim 5, wherein the step of fluorescently examining said DNA fragments includes the steps of:

forming a flow stream of said DNA fragments serially spaced apart in said flow stream;

illuminating said flow stream with a laser effective to cause said first dye to fluoresce; and

measuring the intensity of the total fluorescence from each one of said fragments of DNA in said flow stream.

16. A method according to Claim 15, wherein said step of measuring said total fluorescence intensity further includes the step of differentiating dye molecules bound to said DNA fragments from unbound dye molecules.

17. A method according to Claim 6, wherein the step of fluorescently examining said DNA fragments includes the steps of:

forming a flow stream of said DNA fragments serially spaced apart in said flow stream;

illuminating said flow stream with a laser effective to cause said first dye to fluoresce; and

measuring the total fluorescence intensity from each one of said fragments of DNA in said flow stream.

- 18. A method according to Claim 17, wherein said step of detecting said total fluorescence further includes the step of differentiating dye molecules bound to said DNA fragments from unbound dye molecules.
- 19. A method according to Claim 8, wherein the step of differentiating said dye molecules includes the steps of:

staining said DNA oligonucleotides with a second dye effective for energy transfer to said first dye;

exciting said second dye for energy transfer to said first dye; and

measuring the fluorescence intensity from said first dye to output a signal functionally related to the size of said DNA fragment.

20. A method according to Claim 8, wherein the step of differentiating said dye molecules includes the steps of:

exciting said first dye with a polarized source; and detecting fluorescence from said first dye through a polarization filter oriented in a polarization direction parallel to the polarization of said polarized source.

21. A method according to Claim 8, wherein the step of differentiating said dye molecules includes the steps of:

setting a first detection window related to the lifetime of fluorescence of dye molecules that are not bound to said DNA fragments;

setting a second detection window after said first detection window and related to the lifetime of fluorescence of dye molecules that are bound to said DNA fragments; and

measuring the intensity of the total fluorescence detected within said second window.

22. A method according to Claim 1, further including the steps of:

forming hybridization probes having a predetermined nucleotide sequence with a fluorescent dye tag;

hybridizing said hybridization probes to said DNA fragments; and

fluorescently examining said fluorescent dye tags to determine the association of said probes with fragment size.

- 23. A method according to Claim 1, further including the step of sorting said stained DNA fragments as a function of said fluorescence intensity signal.
- 24. A method according to Claim 1, further including the steps of:

staining said DNA strand or DNA fragments with a third dye that binds to selected oligonucleotides; and

fluorescently examining said DNA fragments for emissions from said third dye to relate said selected oligonucleotide content to fragment size.

24

# Patents Act 1977 Examiner's report to the Comptroller under Section 17 (The Search Report)

Application number

GB 9302411.5

Relevant Technical fields	Search Examiner
(i) UK Cl (Edition L ) C3H (HB3), G1B (BAC, BAX, BBN)	
(ii) Int CI (Edition 5 ) GO1N, C12Q	DR D ELSY
Databases (see over) (i) UK Patent Office	Date of Search
(ii) ONLINE DATABASE: WPI	19 APRIL 1993

Documents considered relevant following a search in respect of claims 1-24

Category (see over)	Identity of docu	ment and relevant passages	Relevant to claim(s)
A	WO 90/10718	(MILLIPORE) see claims	1
A	WO 89/03432	(UNITED STATES) see claims	1
			•
	·		
	·		
			· ·
·			
		·	
<del></del>	<u>i</u>		

Category	Identity of document and relevant passages	Relevant to claim(s
	•	
	•	
	•	
	•	
		·
	•	

#### **Categories of documents**

- X: Document indicating lack of novelty or of inventive step.
- Y: Document indicating lack of inventive step if combined with one or more other documents of the same category.
- A: Document indicating technological background and/or state of the art.
- P: Document published on or after the declared priority date but before the filing date of the present application.
- E: Patent document published on or after, but with priority date earlier than, the filing date of the present application.
- &: Member of the same patent family, corresponding document.

Databases: The UK Patent Office database comprises classified collections of GB, EP, WO and US patent specifications as outlined periodically in the Official Journal (Patents). The on-line databases considered for search are also listed periodically in the Official Journal (Patents).